Heterogeneous versus homogeneous crystal nucleation of hard spheres

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Hard-sphere model systems are well-suited in both experiment and simulations to investigate fundamental aspects of the crystallization of fluids. In experiments on colloidal models of hard-sphere fluids, the fluid is unavoidably in contact with the walls of the sample cell, where heterogeneous crystallization may take place. In this work we use simulations to investigate the competition between homogeneous and heterogeneous crystallization. We report simulations of wall-induced nucleation for different confining walls. Combining the results of these simulations with earlier studies of homogeneous crystallization allows us to assess the competition between homogeneous and heterogeneous nucleation as a function of the wall type, fluid density and the system size. On flat walls, heterogeneous nucleation will typically overwhelm homogeneous nucleation. Even for surfaces randomly coated with spheres with a diameter that was some three times larger than that of the fluid spheres – as has been used in some experiments – heterogeneous nucleation is likely to be dominant for volume fractions smaller than $\approx 0.535$. Only for a disordered coating that has the same structure as the liquid did we find that nucleation was likely to occur in the bulk. Hence, such coatings might be used to suppress heterogeneous nucleation in experiments. Finally, we report the apparent homogeneous nucleation rate taking into account the formation of crystallites both in the bulk and at the walls. We find that the apparent overall nucleation rates coincide with those reported in “homogeneous nucleation” experiments. This suggests that heterogeneous nucleation at the walls could partly explain the large discrepancies found between experimental measurements and simulation estimates of the homogeneous nucleation rate.

I. Introduction

Crystallization is ubiquitous, and a quantitative understanding of this phenomenon is important for all processes, both basic and applied, where crystallization plays a role.

The crystallization of a disordered assembly of building blocks (atoms, molecules, colloids, proteins...) must be initiated by the nucleation of a crystalline embryo somewhere in the system. The critical nucleus may either emerge in the homogeneous bulk, or heterogeneously, on a surface or on an impurity. The latter mechanism is often faster because it reduces the surface free energy and thereby the nucleation barrier of the emerging solid.

In this work we examine the competition between homogeneous and heterogeneous nucleation for a system composed of hard spheres, which is archetypal for the study of the fluid-to-crystal transition. The hard-sphere model was originally generated in computer simulations, but it could, to a good approximation, be realized in experiments on colloidal suspensions. However, in these experiments the sample is unavoidably contained in a cell and therefore crystallization may occur either in the bulk or on the cell walls.

Here, we use computer simulations to assess the competition between homogeneous and heterogeneous crystallization in hard sphere fluids.

As simulation data on homogeneous hard-sphere nucleation are readily available, we focus on the heterogeneous nucleation taking place on the walls confining the hard sphere fluid. The competition between homogeneous and heterogeneous nucleation depends on the system size as the rates of homogeneous and heterogeneous nucleation of clusters are proportional to the volume and the area of the cell, respectively. As a consequence the dominant nucleation mechanism will depend on...
the system size and the volume fraction occupied by the spheres, ϕ. To assess the homogeneous vs. heterogeneous competition we perform estimates of the time required for a crystal to appear via either mechanism. The homogeneous nucleation rate, \( J_{\text{hom}} \), has been previously estimated with a range of different simulation techniques such as umbrella sampling,\(^5,10\) forward-flux sampling,\(^10\) lattice mold,\(^11\) or seeding.\(^12\) Here, we use a reasonably good fit for \( J_{\text{hom}}(\phi) \) provided by seeding simulations.\(^12\) For heterogeneous nucleation on a flat wall, \( J_{\text{het}} \) the rate had been computed previously for a single state point, using umbrella sampling.\(^13\) In the present paper we report calculations of \( J_{\text{het}} \) as a function of \( \phi \) using “brute-force” molecular dynamics simulations. Both flat and coated walls are considered.

We find that, for a given system size, there is a crossover from heterogeneous to homogeneous nucleation as the density increases. The crossover is due to the fact that the homogeneous nucleation rate increases more sharply with density. Moreover, our simulations predict that, for walls that are either flat or randomly coated with spheres about three times larger than the fluid ones, suppression of heterogeneous nucleation is not viable for \( \phi < 0.535 \), unless astronomically large samples are considered.

This raises the question if any wall coating can be designed that would make heterogeneous nucleation less important than homogeneous nucleation. We find that an immobilized wall coating replicating the structure of the fluid could efficiently prevent heterogeneous nucleation. Finally, by combining homogeneous and heterogeneous nucleation rates we compute an apparent homogeneous nucleation rate that takes into account crystallites nucleated both in the bulk and at the cell walls. We find that the apparent rate matches reported experimental data for \( J_{\text{hom}} \).\(^14–16\) Our findings suggest that careful corrections for heterogeneous nucleation are needed when comparing experimental and numerical estimates of the homogeneous nucleation rate.\(^5,10\)

II. Results

A. Heterogeneous nucleation rate on flat walls

The heterogeneous nucleation rate, \( J_{\text{het}} \), is defined as the number of clusters that nucleate per unit time and area:

\[
J_{\text{het}} = \frac{1}{A_{\text{het}}} \tag{1}
\]

where \( A \) is the area of the walls, and \( t_{\text{het}} \) is the average time required for a critical crystal nucleus to appear at the walls.

To obtain \( J_{\text{het}} \) we performed molecular dynamics (MD) simulations in a box with a (quasi) flat hard wall perpendicular to \( z \) and periodic boundary conditions in the other two directions. The size of the flat hard wall is \( 11.5 \times 11.5 \sigma^2 \), where \( \sigma \) denotes the hard sphere diameter.

We performed simulations with 2548 fluid particles. We fixed the pressure with a barostat along the \( z \) direction. To perform MD simulations (with the GROMACS package\(^17\)) we use the pseudo-hard sphere potential,\(^18\) where the discontinuous hard sphere interaction is replaced by the expression:

\[
\begin{align*}
50 \left( \frac{50}{49} \right) \sigma & \quad \text{for} \quad r < \left( \frac{50}{49} \right) \sigma \\
0 & \quad \text{for} \quad r \geq \left( \frac{50}{49} \right) \sigma
\end{align*}
\]

This model mimics the equilibrium\(^18,19\) and dynamic\(^18,20,21\) properties of hard spheres. Approximating hard spheres by very steep but continuous potentials allows us to use a constant time-step MD code, such as the one implemented in GROMACS. These codes have the advantage that they allow for highly efficient parallel simulations, which is important for the large system sizes needed to study nucleation. In contrast, event driven hard-sphere MD requires active load-balancing. Whilst parallel, event-driven MD codes have been proposed (see, e.g. ref. 22) we opted for the GROMACS approach, because it is versatile and has been extensively tested.

In practice, the flat wall is built by positioning spheres with the same size as the fluid ones in a single-layer square lattice of a 0.7\( \sigma \) lattice spacing. This makes a virtually flat slab whose particles interact with the fluid ones via the pseudo-hard sphere potential. We have checked with specific cases that by reducing the lattice spacing of the spheres that form the wall to 0.4\( \sigma \) we obtain the same results. This confirms that the wall is effectively flat already with a 0.7\( \sigma \) spacing. The interaction between the fluid and the wall spheres is given by the pseudo-hard sphere potential defined above. All other simulation details are the same as in our previous publications using the pseudo-hard sphere potential.\(^12,19\)

In Fig. 1 we show a sequence of snapshots illustrating a heterogeneous crystallization event where a crystal cluster nucleates on the wall. In Fig. 2 we plot the time evolution of the number of particles in the largest crystal cluster (detected with a Steinhardt-like order parameter\(^24,25\) as implemented in ref. 12 and 23). The figure shows different trajectories obtained at fixed \( \phi \). Clearly, after an induction period, nucleation takes place.

![Fig. 1](image-url) Sequence of snapshots showing a heterogeneous nucleation event at \( \phi = 0.52 \). The box is filled with particles, but only those detected to belong to the largest crystalline cluster according to a specific local-bond order parameter\(^22,23\) are shown.
place and the number of particles with a crystalline environment (“crystalline particles”) rises sharply. To obtain the average time required to observe one heterogeneous nucleation event, \( t_{\text{het}} \), we use some 8 trajectories for every pressure (each corresponding to a given choice of \( f \)). We then obtain \( J_{\text{het}} \) via eqn (1) (note that the area has to be multiplied by two to take into account that crystals can be formed on either side of the wall). We repeat this procedure for several values of \( f \) and obtain \( J_{\text{het}}(f) \) as shown in Fig. 3 in red. Our values are consistent with those determined by Auer and Frenkel for low \( f \) using umbrella sampling (pink point in the figure). Such consistency validates our simulation approach to obtain heterogeneous nucleation rates of hard spheres on hard flat walls. These results are not strongly sensitive to the interaction between the wall and the fluid particles: if this interaction is changed from pseudo-hard sphere to Lennard-Jones we obtain the same results within the uncertainty of our calculations. This is consistent with the fact that a small softness\(^{26}\) (or an anisotropic shape\(^{27,28}\)) do not change the nucleation rate very much.

**Fig. 3** Red/blue: heterogeneous nucleation rate on a flat/LSC wall as a function of \( \phi \). In pink we include the data obtained by Frenkel and Auer with umbrella sampling for a flat wall.\(^{15}\)

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**B. Homogeneous vs. heterogeneous nucleation**

Once \( J_{\text{het}}(\phi) \) is known, the average time required for a cluster to appear at the walls, \( t_{\text{het}} \), can be obtained via eqn (1). We will compare this time with the one required for a critical cluster to appear in the bulk, \( t_{\text{hom}} \):

\[
t_{\text{hom}} = \frac{1}{V J_{\text{hom}}}
\]

where \( V \) is the volume and \( J_{\text{hom}} \) is the homogeneous nucleation rate. We use a fit for \( J_{\text{hom}} \) obtained in simulations of seeded crystallization and inspired by classical nucleation theory.\(^{12}\)

Such a fit is consistent with independent calculations of the nucleation rate by means of other techniques like umbrella sampling, forward-flux sampling and brute-force simulations.\(^{10,12,29,30}\)

From eqn (1) and (3), it is evident that to estimate \( t_{\text{het}} \) and \( t_{\text{hom}} \) for a given sample-cell geometry, we need to specify the volume and the surface area of the sample cell. Here we consider a cubic cell of \( 1 \times 1 \times 1 \) cm\(^3\) and particles of 200 nm radius, which are typical values for light-scattering experiments.\(^{14-16}\) The resulting \( t_{\text{hom}}(\phi) \) and \( t_{\text{het}}(\phi) \) curves are shown in green and red respectively in Fig. 4. For a given \( \phi \), the dominant mechanism is the one with the smallest nucleation time. Clearly, at low \( \phi \) heterogeneous nucleation is rapidly becoming dominant. The curves cross at \( \phi \approx 0.54 \). This is in fairly good agreement with experimental studies on the competition between homogeneous and heterogeneous crystallization in hard sphere-like colloids.\(^{31}\) Thus, our simulations suggest that, for flat walls and the system size under consideration (\( 1 \times 1 \times 1 \) cm\(^3\) and 200 nm radius particles), bulk crystal formation can only be observed for \( \phi \geq 0.54 \). The reason why there is a crossover between heterogeneous and homogeneous nucleation is that \( J_{\text{hom}} \) varies more sharply with \( \phi \) than \( J_{\text{het}} \). In Fig. 3 it can be seen that \( J_{\text{het}} \) only increases by about 5 orders of magnitude in going from \( \phi = 0.5 \) to \( \phi = 0.54 \). By contrast, \( J_{\text{hom}} \) increases by hundreds of orders of magnitude in the same density interval.\(^{12}\)

We note that since \( t_{\text{hom}} \) and \( t_{\text{het}} \) depend on the volume and the area of the cell, respectively, the crossover \( \phi \), \( \phi_{c} \), is system
size dependent. In Fig. 5 we plot $\phi_c$ versus the length of the edge of a cubic cell for particles of 200 nm radius. The red curve corresponds to a flat wall, which is the case we are discussing so far. The $\phi_c$ curve can be interpreted as a prediction of the minimum system size required to measure homogeneous nucleation without the interference of crystallization at the walls for a given value of $\phi$. Particles with a 200 nm radius are taken to perform this calculation. For any other particle size the x axis should be accordingly rescaled by the factor $r/200$ nm, where $r$ is the radius of interest.

C. Coated walls

Thus far we have considered heterogeneous nucleation on flat walls. An obvious question is to what extent roughening the walls can suppress heterogeneous nucleation. A number of experimental studies on homogeneous nucleation have attempted to suppress heterogeneous nucleation by coating the walls of the sample cell with sintered particles. Such a coating is meant to suppress the layering that is induced in the liquid by a flat wall. Unfortunately, the studies that explicitly mention the use of a coating focus on crystallization of charged rather than hard colloids. For instance, in the work by Ziese et al. the walls are coated with particles that had a diameter 3.4 times larger than those of the fluid particles.

Inspired by this experimental work, we have repeated our heterogeneous nucleation study with a coated wall built by randomly covering a plane with non-overlapping hard spheres 3.4 times larger than the fluid ones (specifically, 142 spheres are located on a square whose side is 14 times the diameter of a single fluid sphere). All spheres have their center on the same plane. We refer to this wall as a “large sphere coating” (LSC).

The blue data in Fig. 3 show that, for such a coating, the heterogeneous nucleation rate goes down by several orders of magnitude. Consequently, $t_{\text{het}}$ increases (in blue in Fig. 4) and the crossover density becomes smaller (in blue in Fig. 5). However, the effect of the coating does not greatly alleviate the problem that unrealistically large samples are required to eliminate the effect of heterogeneous nucleation for volume fractions lower than $\phi = 0.525$, where large discrepancies have been found between experimental and simulation data on homogeneous nucleation rates.

The question that arises now is whether there is an ideal coating that would suppress heterogeneous nucleation in experiments. A intuitive idea would be to use an amorphous solid coating with the same structure as the liquid. We refer to this kind of wall as “frozen-liquid”. It seems plausible that the crystal–frozen-liquid interfacial free energy would be slightly higher than that of the crystal–liquid. The reason is that the frozen-liquid wall, unlike the liquid, is unable to accommodate the capillary fluctuations of the crystal–liquid interface and the local structuring of a liquid close to a crystal interface. Suppressing both effects by an external potential would require reversible work, and hence the crystal–frozen wall interfacial free energy is likely to be higher than that of the crystal–liquid interface. If that is the case, the crystal will not “wet” the frozen wall and therefore heterogeneous nucleation will not happen.

To test this idea we performed simulations at $\phi = 0.535$ with a frozen-liquid wall at the same density as the fluid, as shown in Fig. 6. The system in the figure is composed of 2548 fluid particles and the area of the simulation box side parallel to the solid wall is $11.5 \times 11.5 \sigma^2$. The average crystallization time in such a system over 5 trajectories is $12000 \pm 1000 \sigma^2/(6D_s)$. This time is virtually identical to that obtained with no wall and the same number of fluid particles: $11000 \pm 1000 \sigma^2/(6D_s)$. Moreover, we observe that crystal nuclei typically appear away from the frozen-liquid wall. In addition, when we double the system size, leaving the area of the frozen-liquid wall fixed, the crystallization time is roughly halved. Therefore, it seems that under the conditions studied, the frozen-liquid wall is able to suppress heterogeneous nucleation completely. We recall that on the LSC wall the crystallization time goes down from $O(10^3) \sigma^2/(6D_s)$ to $75 \pm 10 \sigma^2/(6D_s)$ because nucleation takes place heterogeneously. With a flat wall crystal nucleation is even faster ($2 \pm 10 \sigma^2/(6D_s)$). Both for the flat and the LSC walls, the observed crystallization time does not change when doubling the
system volume while keeping the wall area constant. In summary, in the presence of a “frozen-liquid” wall, we were able to reproduce homogeneous nucleation rates at \( \phi = 0.535 \). Therefore, the use of a solid coating with the structure of the fluid could be a promising strategy to suppress heterogeneous nucleation in experiments. However, our study does not rule out the existence of other types of coatings also capable of effectively suppressing heterogeneous nucleation.

Let us finish this section by comparing the structure of the fluid in all types of walls studied: flat, LSC and frozen-liquid. In Fig. 7 we show the fluid density profile along the direction perpendicular to the wall for \( \phi = 0.535 \). For reference, we also include the density profile of the fluid in a box with periodic boundary conditions (no walls). Clearly, the flat wall induces strong layering in the fluid, the LSC wall also induces evident layering, although less pronounced, and the frozen-liquid wall does not induce any clear layering. Given that layering enhances the crystal nucleation ability, the density profiles shown in Fig. 7 help rationalise our results that heterogeneous nucleation is readily induced by flat and LSC walls (by the former more efficiently than by the latter) and that it is prevented by frozen-liquid walls.

D. On the discrepancy between experiments and simulations

Experimental measurements\(^{14–16} \) and numerical calculations\(^ {5,10} \) of \( J_{\text{hom}} \) agree at high densities but differ by many orders of magnitude for lower values of \( \phi \).\(^ {5,10} \) The situation is summarised in Fig. 8, where simulation estimates and experimental measurements of the rate are shown in green and purple respectively. The discrepancy between simulations and experiments is frustrating considering the simplicity of the system. However, on the basis of the heterogeneous nucleation rates that we have computed, we may reconcile experiment and simulation. We define an apparent homogeneous nucleation rate, \( J_{\text{ap}} \), that takes into account the simultaneous contribution of homogeneous and heterogeneous nucleation:

\[
J_{\text{ap}} = \frac{1}{t_{\text{ap}}} V
\]  

(4)

![Fig. 7](image)

**Fig. 7** Fluid density profile along the direction perpendicular to the wall for flat, LSC and frozen-liquid walls for a fluid at \( \phi = 0.535 \). The density profiles start at zero from the middle of wall. The fluid density profile under periodic boundary conditions (no wall) is also included for visual reference (dashed red).

where \( V \) is the volume of the sample and \( t_{\text{ap}} \) is the time required for the first critical cluster to appear, be it homogeneously or heterogeneously. The inverse of \( t_{\text{ap}} \) is the frequency of appearance of a critical cluster in the system. Such a frequency will contain the contribution of two terms, one from homogeneously and another from heterogeneously nucleated crystallites:

\[
\frac{1}{t_{\text{ap}}} = \frac{1}{t_{\text{hom}}} + \frac{1}{t_{\text{het}}}
\]  

(5)

Using the numerical data for \( J_{\text{hom}} \) and \( J_{\text{het}} \) we can estimate \( t_{\text{ap}} \) and hence \( J_{\text{ap}} \). We use for our calculations particles of a 200 nm radius and a 1 \times 1 \times 3 \text{ cm} \text{ cell as in ref. 14. The resulting }\( J_{\text{ap}} \) is shown in Fig. 8 in red for flat walls and in blue for a random coating of larger spheres (size ratio 3.4). At large volume fractions (\( \phi > 0.54 \)) the apparent rate curves coincide with the homogeneous nucleation curve from simulations (dashed green) and with experimental data (purple symbols). This is to be expected because the frequency of appearance of crystallites becomes increasingly dominated by the homogeneous contribution. At lower densities the heterogeneous contribution to the crystallization frequency in eqn (5) causes a departure of the \( J_{\text{ap}} \) curves from the \( J_{\text{hom}} \) simulation curve. The blue curve departs from the dashed green one at lower densities because the coating lowers \( J_{\text{het}} \) (see Fig. 3). In other words, randomly coating the walls with larger spheres widens the density range over which experimental measurements are not affected by heterogeneous nucleation, but only slightly. It is quite suggestive that the red \( J_{\text{ap}} \) curve (flat walls) coincides with the experimental data from ref. 14–16, where the cell walls were presumably flat (at least, these papers do not mention the use of any wall coating). Of course, we cannot rule out that other factors, such as sedimentation, incomplete shear melting, impurities, uncertainties in the determination of the volume
fraction or hydrodynamic effects, also play a role in the experimental determination of the crystallization rate.\textsuperscript{34–36} However, we argue that a quantitative study of these more subtle effects can only be carried out once the experimental data have been properly corrected for wall-induced crystal nucleation. Our results suggest that applying such a correction may go a long way towards reconciling the many orders of magnitude discrepancy between the homogeneous nucleation rate obtained in simulations and the apparent homogeneous nucleation rate observed in experiments.

III. Summary and conclusions

In summary, we have used computer simulations to study the competition between homogeneous and heterogeneous crystal nucleation in hard-sphere suspensions. To estimate the relative importance of homogeneous and heterogeneous nucleation we have performed systematic simulations of the rate of heterogeneous crystal nucleation for a range of different conditions, and combined the results with existing numerical data on homogeneous crystal nucleation.

We find a crossover from heterogeneous to homogeneous nucleation as the density increases. The crossover is due to the fact that the homogeneous nucleation rate increases more sharply with density than the heterogeneous rate. The crossover density depends on the system size and on the shape of the solid surface on which heterogeneous nucleation takes place. We find that, for the wall surfaces that we studied (flat, and randomly coated with spheres about 3 times larger than the suspended colloids), avoiding heterogeneous nucleation for $\phi < \sim 0.535$ is not feasible for reasonable sample sizes. Therefore, we predict that the studied walls would not avoid heterogeneous nucleation in the density range where large discrepancies between experimental measurements and simulation estimates of the homogeneous nucleation rate have been found. We propose an amorphous coating based on the structure of the liquid that could effectively suppress heterogeneous nucleation.

Finally, we show that a homogeneous nucleation rate that takes into account both homogeneously and heterogeneously nucleated crystallites for cell sizes typically used in light scattering experiments coincides with the experimentally reported nucleation rate, provided that the walls are flat. Thus, our work therefore suggests that heterogeneous nucleation at the cell walls may at least partly explain the reported discrepancy of many orders of magnitude between experimental measurements and simulation estimates of the homogeneous nucleation rate.

Conflicts of interest

There are no conflicts to declare.

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